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The Genome of *Cardinium* cBtQ1 Provides Insights into Genome Reduction, Symbiont Motility, and Its Settlement in *Bemisia tabaci*

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Data deposition: *Cardinium* cBtQ1 genome has been deposited at GenBank/EMBL/DDJB databases under the accession CBQZ010000001–CBQZ010000011 for the chromosomal contigs and HG422566 for the plasmid. Cytochrome oxidase I (COI) sequences from *Bemisia tabaci* and 16S rRNA sequences from *Cardinium* sp. derived from this work have been deposited at GenBank/EMBL/DDJB under accession HG421085–HG421096 and HG421077–HG421084, respectively.

Abstract

Many insects harbor inherited bacterial endosymbionts. Although some of them are not strictly essential and are considered facultative, they can be a key to host survival under specific environmental conditions, such as parasitoid attacks, climate changes, or insecticide pressures. The whitefly *Bemisia tabaci* is at the top of the list of organisms inflicting agricultural damage and outbreaks, and changes in its distribution may be associated to global warming. In this work, we have sequenced and analyzed the genome of *Cardinium* cBtQ1, a facultative bacterial endosymbiont of *B. tabaci* and propose that it belongs to a new taxonomic family, which also includes *Candidatus* Amoebophilus asiaticus and *Cardinium* cEper1, endosymbionts of amoeba and wasps, respectively. Reconstruction of their last common ancestors' gene contents revealed an initial massive gene loss from the free-living ancestor. This was followed in *Cardinium* by smaller losses, associated with settlement in arthropods. Some of these losses, affecting cofactor and amino acid biosynthetic encoding genes, took place in *Cardinium* cBtQ1 after its divergence from the *Cardinium* cEper1 lineage and were related to its settlement in the whitefly and its endosymbionts. Furthermore, the *Cardinium* cBtQ1 genome displays a large proportion of transposable elements, which have recently inactivated genes and produced chromosomal rearrangements. The genome also contains a chromosomal duplication and a multicopy plasmid, which harbors several genes putatively associated with gliding motility, as well as two other genes encoding proteins with potential insecticidal activity. As gene amplification is very rare in endosymbionts, an important function of these genes cannot be ruled out.

Key words: Amoebophilaceae, IS elements, gliding motility, *Candidatus* *Cardinium* hertigii, host–symbiont interaction.

Introduction

The sweet potato whitefly *Bemisia tabaci* (Aleyrodidae) is an important polyphagous agricultural pest (Byrne and Bellows 1991). It feeds on plant phloem, causing many economic losses, affecting the plants directly and/or indirectly by transmitting several viruses. *Bemisia tabaci* was originally described

as a complex species comprised by many biotypes (Brown et al. 1995), but recently it has been suggested that it is actually a complex of at least 24 morphologically indistinguishable species (De Barro et al. 2011), although classification in biotypes still persists. Among the most important biotypes are B (Middle East-Asia Minor 1 species), which arose in the

Middle East but now has a cosmopolitan distribution, and Q (Mediterranean species), which originated in the Mediterranean basin and is also widely distributed at present. A phylogenetic analysis of mitochondrial biotype Q individuals worldwide has revealed the existence of three subclades (Q1–Q3) (Gueguen et al. 2010).

Similar to other phloem-feeding insects, which need essential amino acids and other nutrients due to their unbalanced diet, whiteflies have established mutualistic relationships with bacterial endosymbionts that provide essential compounds (Baumann 2005; Moran et al. 2008; Moya et al. 2008). All whiteflies species harbor the obligate, or primary endosymbiont, “*Candidatus Portiera aleyrodidarum*” (Gammaproteobacteria; hereafter *Portiera*) (Thao and Baumann 2004), which is restricted to specialized insect cells called bacteriocytes. In *B. tabaci*, several other bacterial species (secondary or facultative symbionts) may coexist with *Portiera* (Gottlieb et al. 2008). The most frequently observed are *Arsenophonus* sp. and “*Candidatus Hamiltonella defensa*” (hereafter *Hamiltonella*) (both Gammaproteobacteria), which share bacteriocytes with *Portiera*. Less frequently present are *Rickettsia* sp., *Wolbachia* sp. (both Alphaproteobacteria) and “*Candidatus Cardinium* sp.” (Bacteroidetes), detected in bacteriocytes but also in other insect tissues.

Symbiosis between whiteflies and *Portiera* has been shaped through a long-term relationship, mainly characterized by *Portiera* providing essential nutrients (amino acids and carotenoids) to the host, thus compensating for its deficient diet, as revealed by genome sequencing (Santos-Garcia et al. 2012; Sloan and Moran 2012). However, the effects of facultative endosymbionts need to be determined. So far, it has been noted that secondary symbionts in other insects can have different effects, such as reproductive manipulation, nutritional contribution, temperature tolerance, and defence against pathogens and parasitoids (Feldhaar 2011; White et al. 2011).

“*Ca. Cardinium hertigii*” (hereafter *C. hertigii*) was first characterized in *Encarsia* wasps, which are parasitoids of *B. tabaci*, and it was proposed as the species type (Zchori-Fein et al. 2004). However, in recent years, infections with bacteria belonging to the genus *Cardinium* have been detected not only in whiteflies but also in other insects (armored scale, sharpshooters, and *Culicoides* spp.) and other arthropods (mites, ticks, spiders, and copepods). Nowadays, the infection rate in arthropods has been estimated as close to 7% (Zchori-Fein and Perlman 2004; Gruwell et al. 2009; Nakamura et al. 2009). Based on molecular data (16S rRNA and *gyrB* genes) and the presence of microtubule-like complexes, a morphological feature shared by all known *Cardinium*, the genus has been divided into supergroups and strains, following a nomenclature similar to *Wolbachia* endosymbionts, with four described supergroups (A, B, C, and D) (Lo et al. 2002; Nakamura et al. 2009; Edlund et al. 2012). Similarly, in several arthropod taxa, *C. hertigii* has been described as a reproductive manipulator through diverse effects such as feminization,

cytoplasmic incompatibility, and induction of parthenogenesis (White et al. 2011). However, these effects have not been found in other species (e.g., *B. tabaci*), suggesting that *C. hertigii* might also be a mutualistic endosymbiont. This aforesaid claim has been supported by the recently released genome of *Cardinium* cEper1 (endosymbiont of the wasp *Encarsia pergandiella*), which encodes a complete biotin biosynthetic pathway, suggesting a potential role in wasp nutrition (Penz et al. 2012). The 16S rRNA gene sequences from several *Cardinium* endosymbionts of *B. tabaci* reported to date are very similar (>99%) to those of the species type *C. hertigii*, suggesting that they are strains of the same species. Several analyses of secondary endosymbiont coinfections revealed that *B. tabaci* individuals infected only with *C. hertigii* are very unusual, especially for the C1 strain (supergroup A), the most widespread *Cardinium* among whiteflies (Gueguen et al. 2010). Generally, the coexistence with *Hamiltonella* is the most frequently observed, although combinations with other secondary symbionts, such as *Wolbachia* sp. or *Rickettsia* sp., are also possible (Gueguen et al. 2010; Skajjac et al. 2010; Park et al. 2012).

The laboratory strain *B. tabaci* QHC-VLC, belonging to the Q1 subclade, harbors *Cardinium* cBtQ1, which belongs to the C1 strain according to its 16S rRNA gene. This strain coexists within bacteriocytes harboring *Portiera* and *Hamiltonella* and can also be found scattered in different tissues of the whitefly (Gottlieb et al. 2008), similar to other hosts infected by *C. hertigii* (Bigliardi et al. 2006; Kitajima et al. 2007; Nakamura et al. 2009). In contrast, *C. hertigii* strains from different *Encarsia* species have only been detected in the ovaries and accessory cells (Zchori-Fein et al. 2004; Penz et al. 2012).

In this work, the genome of *Cardinium* cBtQ1 endosymbiont of *B. tabaci* QHC-VLC was sequenced and compared with that of the cEper1 strain and with several other Bacteroidetes, including the parasitic amoeba endosymbiont “*Candidatus Amoebophilus asiaticus*” (hereafter referred to as *A. asiaticus*) (Horn et al. 2001; Schmitz-Esser et al. 2010). Overall, all the analysis indicated that the *Cardinium* cBtQ1 genome has undergone changes to facilitate its recent establishment in *B. tabaci*.

Materials and Methods

Genome Assembly and Annotation

Enriched bacterial samples (Harrison et al. 1989) were collected from approximately 40,000 *B. tabaci* strain QHC-VLC adult whiteflies. DNA was extracted with the JetFlex Genomic DNA purification kit following the manufacturer’s instructions (Genomed). DNA was pyrosequenced using Roche 454 GS FLX Titanium single-end (shotgun) and paired-end (3 kb) libraries, and an Illumina HiSeq2000 mate-pair library (5 kb). The genome assembly was complex due to the high number of

repetitive sequences, around 14% of the genome. This includes the presence of long duplications, almost 100% identical, and a large number of IS (Insertion Sequence elements). Duplicated regions were detected by abnormal coverage peaks, the differential genomic context (different presence of IS elements, genes, nonrepetitive intergenic regions), and paired-ends/mate-paired connections. The ends of most contigs were incomplete IS elements of different types. Because the contigs ending in the same IS type usually did not overlap, we expected a slight underestimation of the percentage of repetitive elements of *Cardinium* cBtQ1 genome. See [supplementary materials and methods, Supplementary Material online \(supplementary file S1, Supplementary Material online\)](#) for a detailed description of the assembling and annotation procedure, as well as the software used.

TBLASTX was used to compare the gene content between plasmids of *Cardinium* cBtQ1 and cEper1, and BLAST hits and gene order were plotted with the genoPlotR package (Guy et al. 2010) from R software (R Development Core Team 2011).

Phylogenetic and Phylogenomic Analyses

High-quality sequenced 16S rRNAs (including fragments of more than 900 bp) were downloaded from SILVA (Quast et al. 2013) and National Center for Biotechnology Information (NCBI) databases. ssu-aligner was employed for the alignment, because it takes into account the secondary structure (based on covariance models) of the 16S rRNA genes (Nawrocki 2009). Predefined masking was selected to ensure reproducibility in future alignments, and finally, the alignment was refined with Gblocks (Castresana 2000) allowing 50% in coverage gaps. General time reversible, with estimates of invariant sites and gamma-distribution among-site rate variation (GTR+I+G), was selected as the best model using jModeltest2 (Darriba et al. 2012). Additionally, amino acid sequences for the *gyrB* gene were downloaded from the NCBI database, aligned with MAFFT using the L-INS-i algorithm (Katoh et al. 2002) and refined with Gblocks. ProtTest3 (Darriba et al. 2011) gave the improved general amino-acid replacement matrix, with gamma distributed rates across sites (LG+G) as the best evolutionary model. All accession numbers are supplied in the [supplementary table S1, Supplementary Material online](#).

For phylogenomic reconstructions, 37 orthologous single copy genes related to the translation/transduction machinery and protein folding functions that were present in 61 Bacteroidetes genomes (with the exception of *Candidatus* Sulcia mulleri CARI that lacked 9 of these genes) and a non-Bacteroidetes species, used as outgroup, were selected using the homology search tool from the Microbial Genome Database (Uchiyama 2003) ([supplementary table S1, Supplementary Material online](#)). Encoding proteins were downloaded and aligned with MAFFT v6.717b (L-INS-i) and

concatenated. The selected best model was LG+G+F based on ProtTest3 results.

The first 15 BLASTP hits using the encoded proteins of *Cardinium* cBtQ1 *cgl* gene (*CHV_c0068*) and the *Leishmania major* genes *cgl* (cystathionine gamma-lyase, LmjF35.3230), *cbf* (cystathionine beta-lyase, LmjF32.2640), and *cgs* (cystathionine gamma-synthase, LmjF14.0460) were selected, and their amino acid sequences downloaded. In addition, the first 15 BLASTP hits against Bacteroidetes for the cystathionine beta-lyase protein (*metC*) from *Escherichia coli* str. K-12 (NP_417481.1) were also selected. Amino acid sequences were aligned with MAFFT v6.717b (L-INS-i), and ProtTest3 was used to select the appropriate model for the alignment, in this case, the LG+G model. For plasmid TraG protein, the first 50 BLASTP best hits were downloaded (including *Cardinium* cEper1 and *A. asiaticus*) and aligned with MAFFT v6.717b (L-INS-i). For TraG, the best model selected with ProtTest3 was LG+G+I.

RaxML (Stamatakis 2006) was used to calculate maximum likelihood phylogenetic trees for all the alignments, using optimizations for branch lengths and model parameters, and 1,000 rapid bootstrap replicates. Models were adjusted for each case. In addition, PhyloBayes3 (Lartillot et al. 2009) was used to infer Bayesian posterior distributions for each phylogenetic tree. In each case, evolutionary model was adjusted to the model selected (described above), and three independent chains were run for each alignment. Following Lartillot et al (2009) recommendations, all effective sizes were greater than 200 and maximum discrepancy between chains was less than 0.1. Finally, a majority rule consensus tree was calculated for each alignment.

Genomic Redundancy and Mobile Elements

NUCmer from MUMmer 3 (Kurtz et al. 2004) was used to plot repetitive regions of *A. asiaticus* (AmAs), *Cardinium* cBtQ1, and *Cardinium* cEper1, using each genome as query and subject. Results were filtered and only sequences with at least 95% identity and 500 bp length were used. To estimate the level of redundancy in the genomes, a BLASTN approach using each genome (chromosome plus plasmid concatenated) as query and subject was performed with e-value cutoff of $1e^{-20}$. The BLASTN results were transferred to a spreadsheet, where any alignment with an identity smaller than 95% was removed. Lines were sorted through several steps to identify the single copy segments of the genome. The repetitive fraction was estimated subtracting the summation of the single copy segments from the total genome length. IS elements were detected as described previously (Gil et al. 2008), refined using the web server ISSaga (Varani et al. 2011) and deposited in ISfinder database. Reference copies for each mobile element were used to search with BLASTX against the nonredundant NCBI database ($1e^{-3}$ e-value cutoff). The 25 best hits for each mobile element were used as the input for MEGAN4

(Huson et al. 2011), and taxonomical assignments were done with the default LCA parameters. The genomes of *A. asiaticus*, *Cardinium* cEper1, and *Cardinium* cBtQ1 (with contigs concatenated in decreasing order, excluding the plasmid) were used to compute nucleotide synteny blocks with progressive Mauve aligner (Darling et al. 2010). *Cardinium* cBtQ1 was set as the reference for gene order and the alignment was plotted with the genoPlotR package.

Orthologous Gene Identification

After phylogenomic reconstruction, all genomes belonging to the order Cytophagales, including both *Cardinium* strains (cBtQ1 and cEper1) and *A. asiaticus* (AmAs), were used for orthologous gene identification. *Flavobacterium johnsoniae* (Bacteroidetes: Flavobacteriales), whose gliding motility has been broadly studied, was selected as the outgroup for the order Cytophagales. All proteins, including those from plasmids, were used as the input for OrthoMCL (Li et al. 2003). OrthoMCL and COG (Clusters of Orthologous Groups of proteins) (Tatusov et al. 2000) profile assignment pipelines were run as described previously (1.5 as inflation value, 70% of match cutoff, and an e-value cutoff of $1e^{-5}$) (Manzano-Marín et al. 2012). Gene clusters may contain zero, one, two, or more genes in each genome. Some gene clusters were manually refined because OrthoMCL failed to recognize some orthologous genes in endosymbionts.

Orthologous genes for *A. asiaticus* and both *Cardinium* strains were classified as core genome (genes shared by the three genomes), genes shared by two of the three organisms and strain-specific genes (supplementary table S2, Supplementary Material online). Euler diagrams were obtained with gplots package (Warnes et al. 2013) from R.

Last Common Ancestor Reconstruction

All genomes used in the OrthoMCL clustering method were used to reconstruct the putative last common ancestor (LCA) gene contents for each node in the phylogeny (fig. 1 and supplementary table S3, Supplementary Material online). The MPR (most parsimonious reconstruction) function in ape package (Paradis et al. 2004) from R was used to infer the ancestral state for each character (gene clusters) in each node (LCA). Pseudogenes for all genomes were manually revised. For orthologous cluster assignments of the pseudogenes, TBLASTX was used (e value of $1e^{-5}$ and an overlap of 80% query subject) against the proteins present in the orthologous clusters. Pseudogenes that did not modify the LCA reconstruction (strain-specific genes) were not considered. Pseudogenes that were mobile elements were also excluded. Parsimony reconstruction for orthologous groups that included the previously selected pseudogenes were checked using parsimony reconstruction of discrete characters in Mesquite (Maddison WP and Maddison DR 2011).

For each reconstructed LCA and genome, COG categories were assigned for each orthologous cluster based on the initial OrthoMCL results and the COG assignment described above (Manzano-Marín et al. 2012). For each orthologous cluster, COG categories with less than a 10% of a cluster, as well as the unassigned category, were removed. The LCA, indeterminations (the presence/absence of the gene in the LCA node could not be determined) were counted as half (0.5), instead of presence (1) or absence (0). Relative percentages of each COG type using LCA4 or LCA2 as reference were plotted using the gplots heatmap2 function without dendrograms and reordering functions. Euler diagram was plotted using gplots. COG profiles, stated as the absolute number of COG categories divided by the total number of COG for each genome or LCA, were plotted as a heatmap with gplots allowing hierarchical clustering (dendrograms are grouping the most similar rows or columns together). Cyclobacteraceae family habitats were obtained from GOLD database (supplementary table S3, Supplementary Material online).

Analyses of *B. tabaci* and *Encarsia* spp. Samples

See supplementary materials and methods and tables S4–S6, Supplementary Material online (supplementary file S1, Supplementary Material online).

Fluorescent In Situ Hybridization

See supplementary materials and methods, Supplementary Material online (supplementary file S1, Supplementary Material online).

Results

General Features of *Cardinium* cBtQ1 Genome

Cardinium cBtQ1 has a relatively small genome (1.065 Mb) composed of a chromosome (1.013 Mb) and a circular plasmid (52 kb) named pcBtQ1 (table 1). The chromosomal sequence is composed of 11 contigs (612, 80.4, 77.6, 73.8, 66.9, 41.4, 30.3, 13.5, 7.4, 5.1, and 4.1 kb, respectively) with average coverages of 90× and 547× for 454 and Illumina platforms, respectively, whereas the plasmid is in a single contig with coverages of 595× (454) and 4,046× (Illumina). Paired-end and mate-pair information confirmed that the plasmid was a single closed circular contig. Also, the higher coverage of the plasmid compared with the chromosomal contigs is indicative of a multicopy plasmid, probably between six and seven copies relative to the chromosome.

We found 709 and 30 coding genes on the chromosome and plasmid, respectively. Many of them were annotated as encoding conserved or hypothetical proteins. We also annotated 156 pseudogenes in the chromosome, 132 derived from transposases and 24 from nontransposase genes (supplementary table S7, Supplementary Material online), and 4 in the plasmid (3 transposases and 1 resolvase). The genome

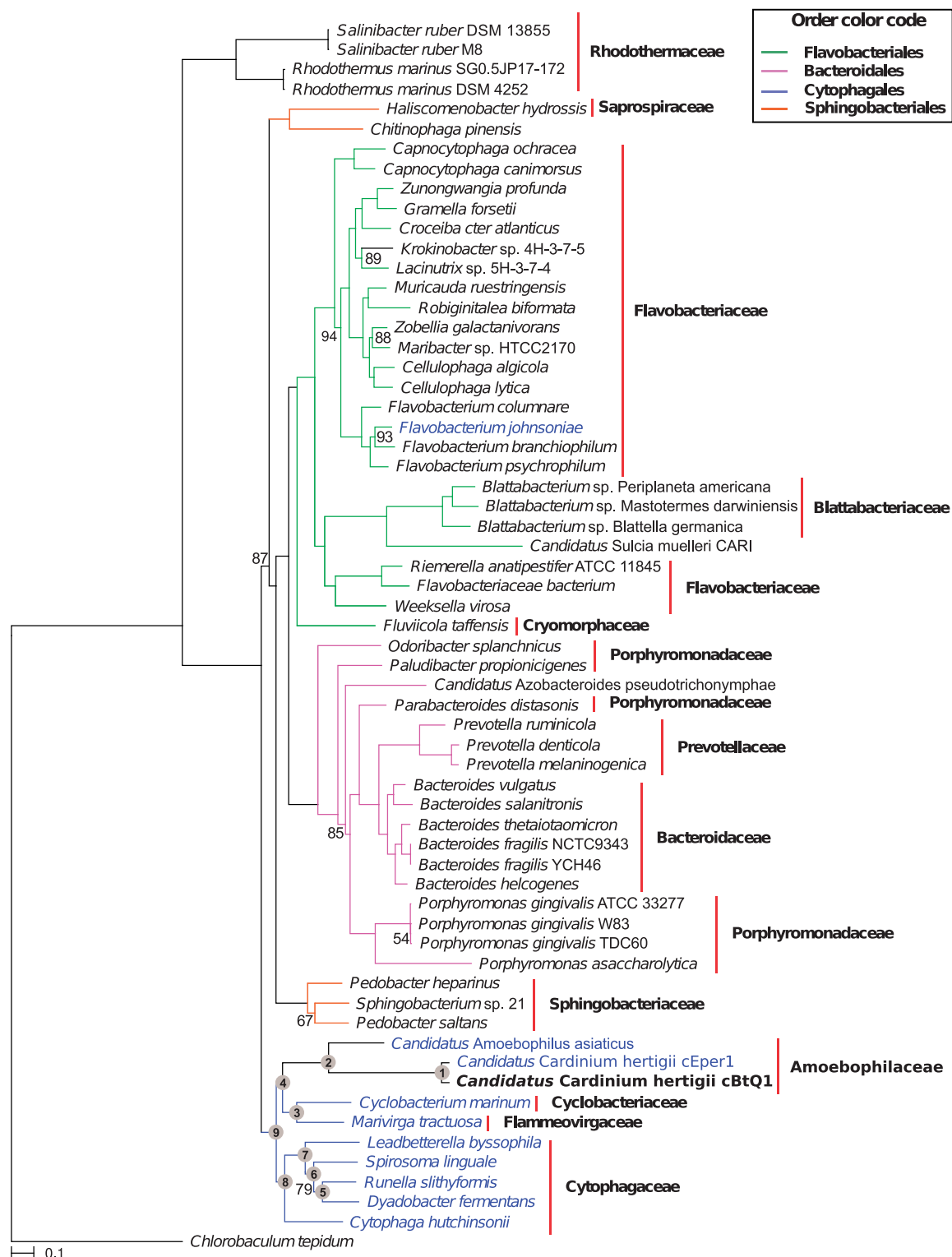


FIG. 1.—Phylogenomic maximum likelihood tree of 61 Bacteroidetes. Phylogenomic reconstruction was done under the LG+G+F model on a concatenated alignment of 37 proteins. *Cardinium* genomes fall in the Cytophagales clade, with *Marivirga tractuosa* and *Cy. marinum* as the closest free-living relatives. *Cardinium* cBtQ1 is displayed in bold. Family names are displayed on the right delimited by a horizontal red line. The genomes used for the LCA reconstruction are shown in blue. Numbers inside gray dots show the LCAs reconstructed in each node. Only maximum likelihood bootstrap values below 95% are displayed. Bayesian posterior probabilities for each node were above 0.95 and are also not displayed. *Chlorobaculum tepidum* was used as outgroup.

Table 1General Genomic Features of *Cardinium* Strains and *Amoebophilus asiaticus*

Bacterial Genome	<i>Cardinium</i> cBtQ1 ^a		<i>Cardinium</i> cEper1 ^b		<i>A. asiaticus</i> 5a2
Host	<i>Bemisia tabaci</i>		<i>Encarsia pergandiella</i>		<i>Acanthamoeba</i> spp.
	Chromosome	Plasmid	Chromosome	Plasmid	Chromosome
Contigs	11	1	1	1	1
Size (kb)	1,013	52	887	58	1,884
GC (%)	35	32	36	31	35
CDS	709	30	841	65	1,557
Average CDS length (bp)	1,033	1,389	911	733	990
Coding density (%)	79.7	80.1	85.5	82.1	81.8
rRNAs	3	—	3	—	3
tRNAs	35	—	37	—	35
Other RNA genes	2	—	—	—	—
Pseudogenes (total)	156	4	3	—	222
Pseudogenes (transposase)	132	3	3	—	—
Pseudogene (other CDS)	24	1	—	—	—
Reference	This study		Penz et al. (2012)		Schmitz-Esser et al. (2010)

^aThe chromosome of *Cardinium* cBtQ1 is not closed, but it is considered a high-quality draft genome.^bThe chromosome of *Cardinium* cEper1 contains a single gap not closed due to repetitive elements.

contains one set of rRNA genes distributed in two segments, one including the 16S rRNA and the other the 23S and the 5S rRNA genes. It also contains a set of 35 tRNA genes, which are able to completely decode the mRNA sequences, and two other noncoding RNA genes (*rnpB* and *tmRNA*) (table 1).

Taxonomic Status of *Cardinium* cBtQ1 Endosymbiont of *B. tabaci* Biotype Q

We performed a phylogenomic reconstruction of a Bacteroidetes phylogeny (see [supplementary table S1, Supplementary Material](#) online, for locus tags in each genome), which showed that both *Cardinium* and *A. asiaticus* formed a well-differentiated clade, related to the families Cyclobacteriaceae and Flammeovirgaceae, with the family Cytophagaceae slightly more distant (fig. 1). This reconstruction was used to select the genomes for subsequent analyses.

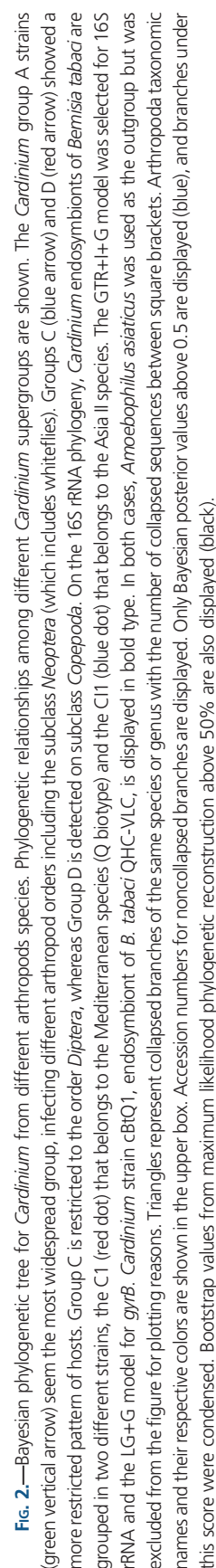
Because this phylogenomic reconstruction is consistent with other reported studies (Gupta and Lorenzini 2007; Karlsson et al. 2011), and due to the high bootstrap values obtained, we propose that the *Cardinium*/*Amoebophilus* clade should be assigned to the order Cytophagales, instead of remaining in the nonclassified Bacteroidetes. Furthermore, the analysis suggests that within the Cytophagales, *Cardinium* forms a new family phylogenetically allied with the Cyclobacteriaceae and Flammeovirgaceae, to which we propose the name Amoebophilaceae (fig. 1).

To establish the relationship of *Cardinium* cBtQ1 to other *Cardinium* endosymbionts based on 16S rRNA sequences, a covariance model aligner was employed. The 16S rRNA alignment was used to infer a phylogeny, showing that almost all *Cardinium* endosymbionts of *B. tabaci* (including cBtQ1) were present in a clade with other *Cardinium* endosymbionts of

several *Encarsia* species (99.14% identity with the whole 16SrRNA of *Cardinium* from *E. pergandiella*) (fig. 2, left). A phylogeny with *gyrB* was also performed, which corroborated the close phylogenetic relation with *Cardinium* from *Encarsia* spp. and also showed that *Cardinium* cBtQ1 was embedded in the *Cardinium*–*Encarsia* clade (fig. 2, right). Because 16S rRNA sequences of *Cardinium* cBtQ1 and the species type *C. hertigii* (symbiont of *E. hispida*) show only 1.2% of differences, we propose that *Cardinium* cBtQ1 is a strain of the latter, in agreement with previous reports (Zchori-Fein and Perlman 2004). The *Bemisia*/*Encarsia* clade belongs to the *Cardinium* group A, which is well differentiated from the other two groups included in the analysis: Group C, which is specific to the genus *Culicoides* (Nakamura et al. 2009) and group D, which is present in some *Copepoda* spp. (Edlund et al. 2012) (fig. 2).

Genome Comparison of *Cardinium* Strains and *A. asiaticus*

Compared with *A. asiaticus* (Schmitz-Esser et al. 2010), the genomes of *Cardinium* cEper1 (Penz et al. 2012) and *Cardinium* cBtQ1 (this work) contain a smaller number of coding genes (table 1). However, *Cardinium* cEper1 has almost no annotated pseudogenes, a difference that might reflect gene annotation criteria because open reading frames that belong to transposase fragments are annotated as coding sequences (CDSs). The average gene identity between *Cardinium* cEper1 and cBtQ1 was 92.9%, whereas the average protein identity was 91.8%. The genome fraction assigned to coding genes (labeled as coding density in table 1) was approximately 6% smaller in *Cardinium* cBtQ1 than in *Cardinium* cEper1.



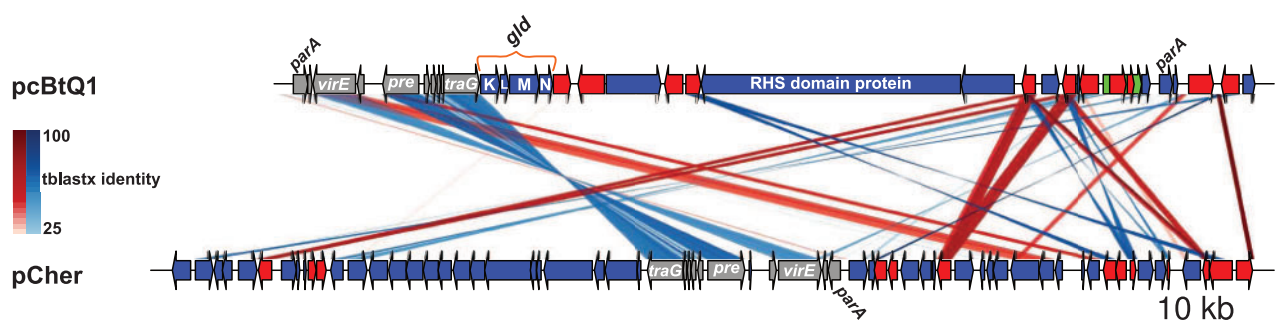


Fig. 3.—Comparison of *Cardinium* plasmids. TBLASTX comparison of the plasmids from *Cardinium* cBtQ1 (pcBtQ1) and cEper1 (pCher). Gray arrows are genes included in the syntenic block, blue arrows nontransposase genes, red arrows transposase genes, and the green arrow is a resolvase pseudogene. Red lines show genes in the same orientation and blue lines in reverse orientation. Some gene names are shown in the plot.

Cardinium cEper1 also contains a plasmid (pCher) of similar size to pcBtQ1. However, the gene content of both plasmids is different, with only a few shared genes in a syntenic segment showing a high level of nucleotide identity (fig. 3). For example, *CHV_p004* (*virE*, virulence-associated E family protein), *CHV_p006* (*pre*, plasmid recombination enzyme), and *CHV_p011* (*traG*, putative conjugal transfer protein TraG) all show nucleotide identities that range from 76% to 83% between both plasmids. Phylogenetic reconstructions for each of these genes support the close relation between both plasmids (see TraG phylogeny in [supplementary file S1: supplementary fig. S1, Supplementary Material](#) online). Nucleotide, gene order conservation, and phylogeny suggest that both plasmids derive from a common ancestral plasmid present before the split of both *Cardinium* lineages. A putative *traG* gene (*Aasi_0886*), closely related to *Cardinium* *traG* phylogenetically ([supplementary fig. S1, Supplementary Material](#) online), is harbored in the *A. asiaticus* chromosome, suggesting that the origin of the plasmid can be traced to the family Amoebophilaceae, with its subsequent chromosomal insertion in *A. asiaticus*.

All three genomes share a core of 468 gene clusters ([supplementary table S2, Supplementary Material](#) online, and [fig. 4](#)), of which six encode putative host interacting proteins (Penz et al. 2012). There are 140 unique gene clusters that were present in both *Cardinium* but not in *A. asiaticus*, 46 of which encode hypothetical proteins, 15 are membrane transport related proteins, and 13 are putative host interacting proteins. Among the remaining *Cardinium* shared genes, there are six coding for transposases, two for phage-derived proteins (or *afp*-like proteins), and five for vitamin biosynthetic proteins ([supplementary table S2, Supplementary Material](#) online).

Cardinium cEper1 has 202 strain-specific gene clusters, which include, among others, those encoding hypothetical proteins (145), transposases (30), host-interacting proteins (6), and biotin (2) and pyridoxal (1) biosynthetic enzymes. *Cardinium* cEper1 and *A. asiaticus* share 13 gene clusters

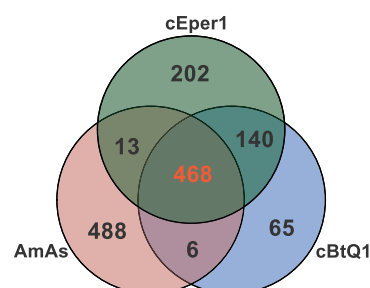


Fig. 4.—Euler diagram of orthologous clusters. Euler diagram representing the core genome, the strain-specific orthologous clusters, and the orthologous clusters shared by only two organisms. Numbers inside each subspace represent the number of orthologous gene clusters assigned to the subspace. Core genome set is displayed in orange. cBtQ1, *Cardinium* cBtQ1; cEper1, *Cardinium* cEper1; AmAs, *Amoebophilus asiaticus*.

with most of them defined as hypothetical proteins (6), mobile elements (3), a cell-wall-related protein, a membrane protein, and a host-manipulation protein ([supplementary table S2, Supplementary Material](#) online).

Cardinium cBtQ1 contains 71 gene clusters, 65 strain-specific, and 6 shared with *A. asiaticus*, which are not present in *Cardinium* cEper1. These gene clusters include ankyrin-domain-containing proteins (14), hypothetical proteins (35), transposases, and other mobile elements (4). A set of very interesting genes is located in the multicopy plasmid of *Cardinium* cBtQ1. They include four gliding genes (*gldK*, *gldL*, *gldM*, and *gldN*, see [fig. 3](#)) that are related to motility in members of the phylum Bacteroidetes (shared with *A. asiaticus*). The fact that the chromosome of *Cardinium* cBtQ1 contains four duplicated genes *rtxBDE* (*A. asiaticus* contained a single copy of the paralogous genes *rtxBE*) and *tolC* related to type I secretion system (T1SS) is also remarkable because only a few sequenced Bacteroidetes harbor secretion system types I, III, IV, or VI (McBride and Zhu 2013). The *rtxBDE* genes, related to the hemolysin secretion proteins (Hly), seem to be

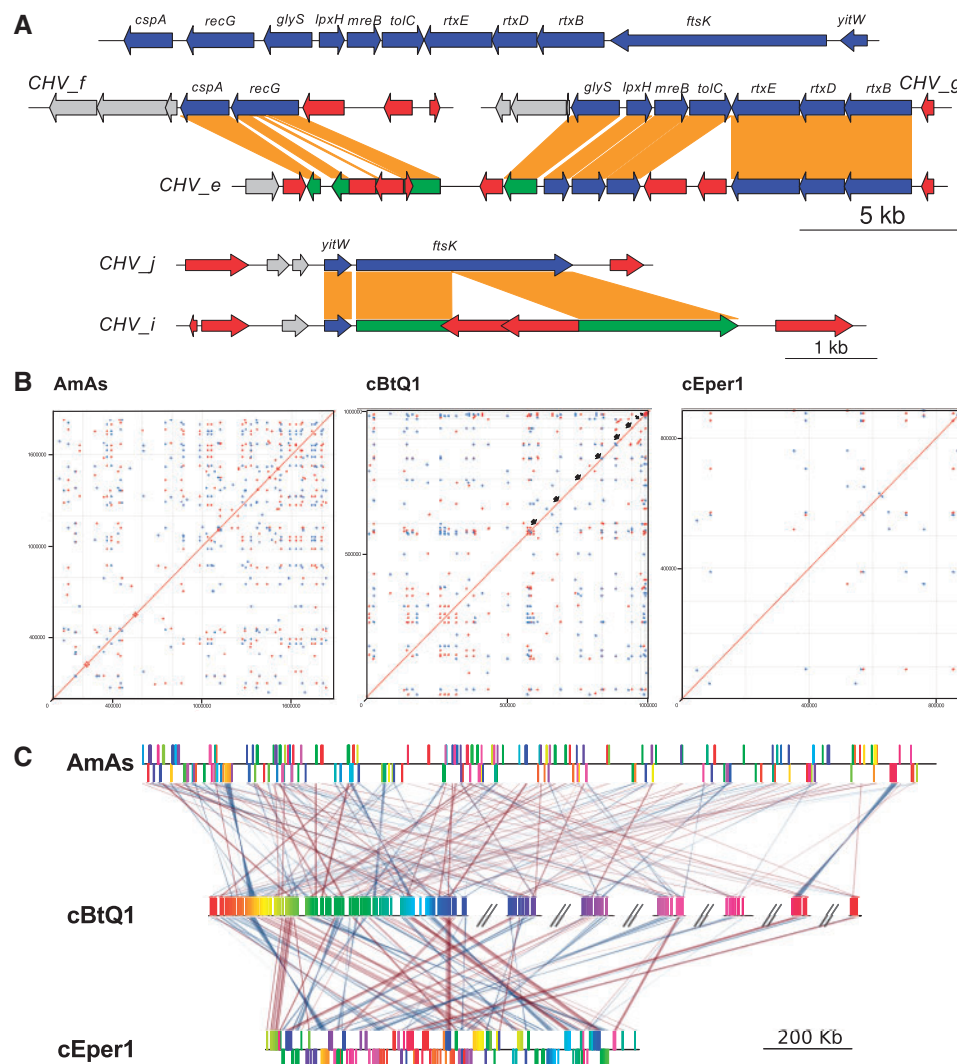


FIG. 5.—Redundancy and synteny between *Cardinium* and *Amoebophilus asiaticus* genomes. (A) Putative linear representation of the ancestral genomic region before duplication (on top) and the present state of the two duplications, which are distributed in five contigs (on bottom). Note that *lpxH*, *mreB*, *tolC*, *rtxBDE*, and *yitW* conserve two intact copies, whereas the other duplicated genes only maintain one intact copy. Red arrows are mobile elements, blue arrows genes in the duplicated region, green arrows pseudogenized genes, and gray arrows adjacent genes outside the duplication. Orange bars connect the two duplicated copies of each gene. Contig names are plotted at the beginning or the end of the contig (*CHV_*), and only regions that contain the duplications are shown. The right ends of contigs *CHV_g* and *CHV_e* are connected through paired-end information with the right ends of either contig *CHV_j* or *CHV_i*. In both cases, a complete ISCa1 copy, whose fragments are detected at the end of the contigs, is required for joining. (B) Mummer plot showing direct (red) and inverted (blue) genomic repeats with at least 500 bp lengths and 95% similarity. For *A. asiaticus* (*AmAs*) and *Cardinium* cEper1 (*cEper1*), inner plot lines denote the division of the chromosome in base pairs sections. Black arrows point contig ends for the largest contigs in *Cardinium* cBtQ1. These contigs were placed in order of decreasing length. Because plots are not scaled to genome size due to limitations of the software, it is noteworthy that the *A. asiaticus* genome is less repetitive than *Cardinium* cBtQ1 although the more compact plot in the former may alter that impression. (C) Common pairwise syntenic blocks of more than 1 kb for *A. asiaticus* (*AmAs*), *Cardinium* cBtQ1 (*cBtQ1*), and cEper1 (*cEper1*). The chromosome of cBtQ1 was taken as reference. Contigs in cBtQ1 are ordered in order of decreasing length and denoted by double backslashes. For plotting reasons, only the seven longest cBtQ1 contigs are shown. Red and blue lines show blocks in direct and inverted orientation. The stronger the line, the more nucleotide identity between syntenic blocks.

an event of horizontal gene transfer (HGT), with RTX toxin transport system of *Vibrio* as best BLAST hits. The chromosomal segment involving these genes is duplicated in *Cardinium* cBtQ1 (fig. 5A).

Other interesting *Cardinium* cBtQ1-specific genes, also located in the plasmid, are the putative toxin-related genes *CHV_p018* and *CHV_p021*. The latter encodes a long (4,603 amino acids) RHS-repeat-associated core-domain protein with

C-terminus ankyrin repeats. Large proteins with RHS domains have been related with bacterial insecticidal toxins and inter-cellular signaling proteins (TIGR03696). Although no clear signal peptide was detected in *CHV_p021*, the presence of ankyrins in the C-terminus domain in combination with a signal peptide has been attributed to protein secretion by T1SS (Kaur et al. 2012). Because the best BLASTX hits, with 63% query coverage and 36% identity (on average), belong to *Daphnia*, *Wolbachia* (HGT event), and mosquitoes, it suggests that the target of this protein can be a conserved protein in arthropods.

The level of redundancy in the genome of *Cardinium* cBtQ1 (~14%) was twice as high as the level found in *Cardinium* cEper1 and *A. asiaticus* (~7% in both cases), most of which is associated with mobile elements (fig. 5B). The mobile elements of *Cardinium* cBtQ1, and their inactive derivatives, account for approximately 166 kb of the chromosome (196 copies) and 12.5 kb of the plasmid (12 copies) (supplementary table S8, Supplementary Material online). From this number of mobile element copies, only 48 contained a functional transposase gene, whereas 135 were transposase pseudogenes. These transposase proteins were classified in 20 different IS families, with only eight being complete IS elements (containing intact transposase genes and inverted repeats at their ends) and could be named according to the ISfinder recommendations and deposited under the names ISCca1-8 (supplementary table S8, Supplementary Material online). Only three mobile element types were specific of the *Cardinium* cBtQ1 (ISCca6, nv_IS3, and the Retron type one), whereas the rest of transposases were shared with *A. asiaticus*, *Cardinium* cEper1 or both.

It would appear that at least some IS in *Cardinium* are still active, in contrast with *A. asiaticus* (Schmitz-Esser et al. 2011), given we can observe very recent gene duplication events (based on > 99.9% nucleotide identity), with one of the copies interrupted by the insertion of an IS (e.g., pseudogenes *recG* or *ftsK*) (fig. 5A). Another important feature is the presence of a repetitive element composed of a copy of ISCca4 and a copy of nv_IS2, resulting in an IS that is apparently active. The inactivation of *ftsK* was produced by the insertion of this chimeric IS. Recombination associated to IS may be the cause of the high number of rearrangements in the *Cardinium* cBtQ1, with only some microsyntenic blocks maintained (fig. 5C).

Finally, *Cardinium* cBtQ1 contains a recent chromosome segmental duplication (almost 100% identical) involving at least 11 genes and around 17 kb, distributed in five contigs (fig. 5A). The duplicated region contains the genes *cpsA* (a protease), *recG* (recombination and DNA repair), *glyS* (aminoacyl-tRNA-ligase), *lpxH* (lipid A biosynthesis), *mreB* (actin-like bacterial cytoskeleton), *tolC* (T1SS transmembrane transporter), *rtxB*, *rtxD*, and *rtxE* (T1SS ABC transporters HlyB and HlyD), *ftsK* (cell cycle and chromosome partitioning), and *yitW* (putative chromosome partitioning related function). Although one of the two copies of *cpsA*, *recG*, *glyS*, and

ftsK is pseudogenized by IS insertions, the rest of the genes conserve the two functional copies (fig. 5A), indicating that their retention could be advantageous for the organism fitness. However, the fact that the two copies are still active due to their recent duplication cannot be ruled out.

Evolution of Gene Repertoires in Lineages of *Amoebophilus* and *Cardinium*

Gene clusters obtained with OrthoMCL were classified according to COG categories and used to reconstruct by maximum parsimony the gene cluster content in the nine LCA corresponding to each node denoted with gray circles (fig. 1). Several analyses were performed to compare the gene content of the present and reconstructed ancestral genomes.

First, hierarchical clustering based on the relative abundance (percentage) of each COG category in each genome was performed (fig. 6). Three main clusters were observed: One that contained the endosymbiotic genomes and the LCA1 and 2; a second that grouped *Marivirgia tractuosa* and *Cyclobacterium marinum* with the LCA3, 4, 8, and 9; and a third that contained the rest of the genomes and LCAs. The second cluster (fig. 6, blue) showed a clear reduction in some COG groups as G and K but an enrichment in the H and J groups when it was compared with the third cluster (fig. 6, yellow). Also, hierarchical clustering grouped both *Cardinium* strains with *A. asiaticus* and LCA1 and LCA2. They showed a stronger conservation of genes in some COG categories such as J and O, when they were compared with the free-living Bacteroidetes, a signal also observed in other symbiotic reduced genomes. LCA4, the ancestor of the *Cardinium*/*A. asiaticus* lineage and family Cyclobacteriaceae, was close to the free-living Cyclobacteriaceae (see habitats in supplementary table S3, Supplementary Material online), suggesting a similar free-living style. Parsimony reconstruction assigned 1,301 gene clusters to LCA4 and the equally parsimonious presence/absence of other 684 gene clusters.

Second, the transition from LCA4 to LCA2 was examined comparing the percentages of gene clusters in each COG category regarding LCA4 (100%) (fig. 7A). It had a strong impact in the number of gene clusters with more than half of them being lost (LCA2, 655 gene clusters plus 36 present/absent) (supplementary table S3, Supplementary Material online). Relatively to LCA4 (100%), there was a high reduction in all categories except for some housekeeping categories such as J, L, and D with more than 80% of gene clusters conserved. Biosynthetic capabilities of LCA2 were clearly reduced (e.g., COG categories C, E, F, and H).

Third, the transition from LCA2 to LCA1 and to both *Cardinium* and *A. asiaticus* was examined comparing the percentages of gene clusters in each COG category regarding LCA2 (100%) (fig. 7B). The transition to LCA1 (649 gene

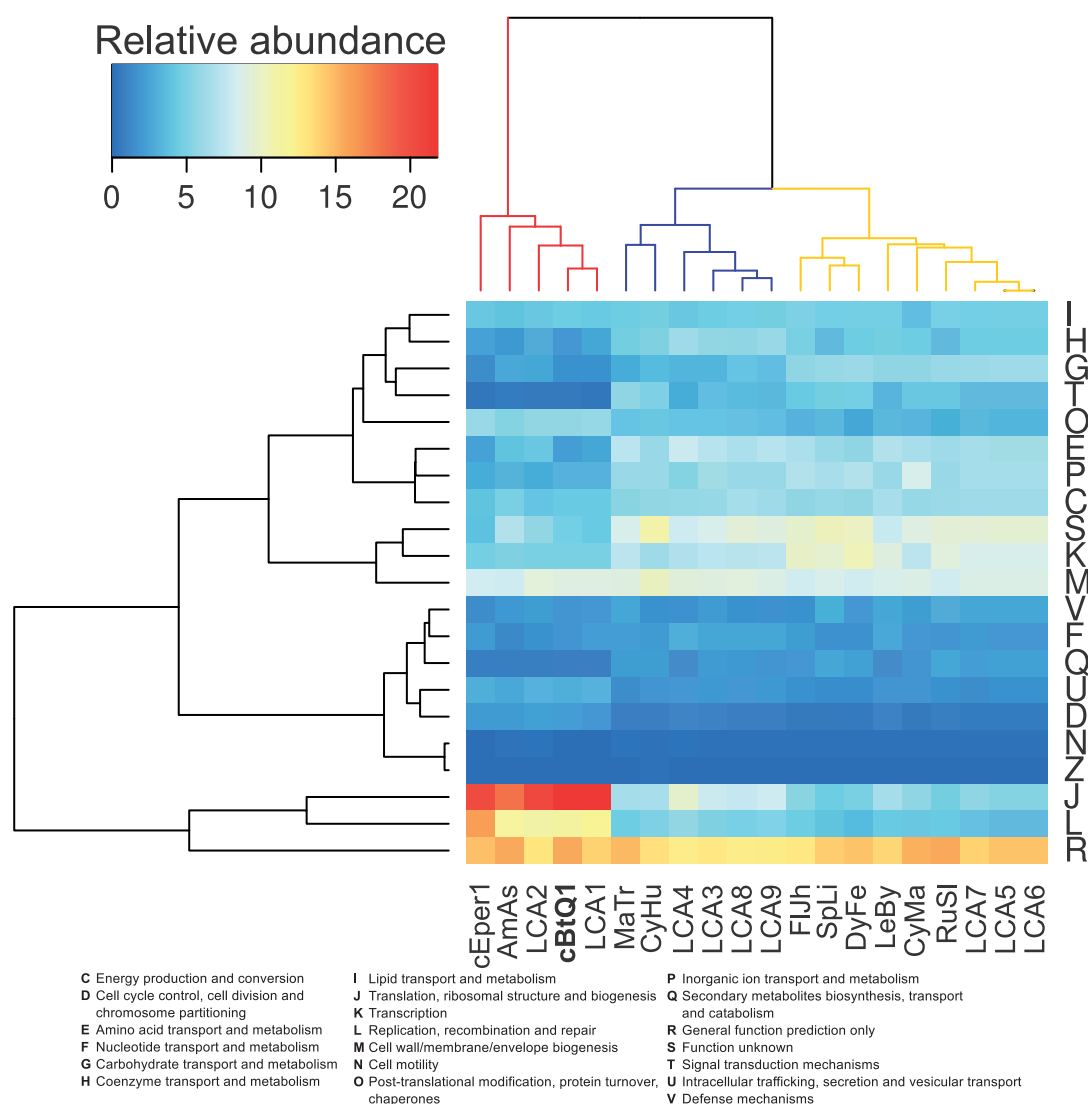


Fig. 6.—Relative abundance of gene clusters in Bacteroidetes and LCAs. Hierarchical clustering heatmap representing the relative abundance (percentage) of each COG category in relation to the total number of gene clusters in each genome. Three main COG clusters (left) are observed: Highly retained categories (J, L, and R), medium retained categories (I, H, G, T, O, E, P, C, S, K, and M), and low retained categories (V, F, Q, U, D, N, and Z). Three main species/LCA cluster (up) are cEper1, AmAs, cBtQ1, LCA1, and LCA2 (only symbionts, left cluster); MaTr, CyHu, LCA3, LCA4, LCA8, and LCA9 (middle cluster), and FIJh, SpLi, DyFe, LeBy, CyMa, RuSl, LCA5, LCA6, and LCA7. Species clustering together by COG categories could have similar metabolic features and consequently, a similar ecological niche. *Cardinium* cEper1 (cEper1), *Ca. Amoebophilus* (AmAs), *Cardinium* cBtQ1 (cBtQ1), *Marivirga tractuosa* (MaTr), *Cytophaga hutchinsonii* (CyHu), *Flavobacterium johnsoniae* (FIJh), *Spirosoma linguale* (SpLi), *Dyadobacter fermentans* (DyFe), *Leadbetterella byssophila* (LeBy), *Cyclobacterium marinum* (CyMa), and *Runella slithyformis* (RuSl).

clusters) produced the loss of 160 gene clusters, although 118 new gene clusters were acquired. Comparing the number of gene clusters of LCA2 to LCA1, and to both *Cardinium* and *A. asiaticus*, we observed several differences among COG categories (fig. 7B, supplementary table S3, Supplementary Material online). First, *A. asiaticus* showed 331 strain-specific gene clusters, distributed in several categories, not present in LCA2. Second, the reductive evolution of the *Cardinium* lineage was more clearly observed in several COG categories, such as E, G, H, S, T, and V. The absence of gene clusters in

Cardinium for the N category was probably due to the fact that some genes related with motility have not been yet annotated in the COG database, especially those involved in gliding motility (see later) that are, in fact, present in *Cardinium* cBtQ1, *A. asiaticus*, LCA1, LCA2, and LCA4.

Biosynthetic Capabilities in *Cardinium* cBtQ1

Cardinium cBtQ1, according to KEGG classification pathways, presents low biosynthetic capabilities, similar to those

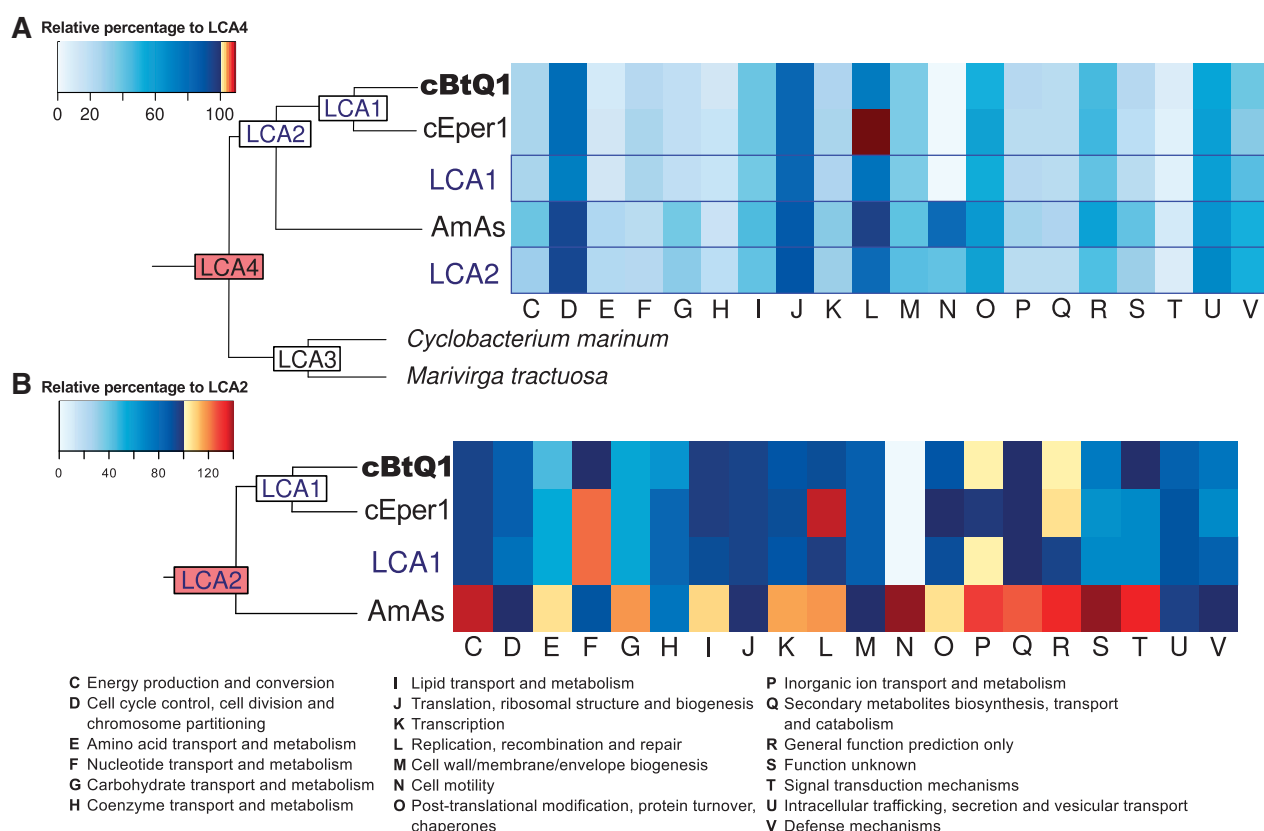


FIG. 7.—Last free-living common ancestor comparison. (A) Heatmap showing the percentage of genes in each COG category, compared with the number of the same category in LCA4 (100%). In left, reduced phylogenomic reconstruction with the name of each LCA reconstructed. (B) The same heatmap type comparing to LCA2 (100%). L category in *Cardinium* cEper1 is an artifact produced by an incorrect annotation of inactivated transposases as CDS instead of pseudogenes. COG definitions are below. cBtQ1, *Cardinium* cBtQ1; cEper1, *Cardinium* cEper1; AmAs, *Amoebophilus asiaticus*.

observed in *Cardinium* cEper1 and *A. asiaticus* (Karlsson et al. 2011; Penz et al. 2012). The main differences between the biosynthetic capabilities of both *Cardinium* strains are in the biosynthesis of vitamins and cofactors. Both bacteria are able to produce lipoate, a key cofactor for intermediate metabolism and an important antioxidant molecule (Spalding and Prigge 2010). *Cardinium* cEper1 has the genes *pdxS* and *pdxT* and can synthesize pyridoxal 5-phosphate (precursor of vitamin B6). However, *pdxT* was pseudogenized by an IS transposition in cBtQ1. This event seems to have happened recently, because the *pdxT* pseudogene is 93.5% identical to the cEper1 gene, a percentage higher than the average gene identity between these two strains.

In addition, it is noteworthy that in *Cardinium* cEper1, the presence of a complete biotin operon, a coenzyme belonging to vitamin B class, is an event of HGT from Alphaproteobacteria to the genus *Cardinium* (Penz et al. 2012). The loss of the ability to synthesize biotin in *Cardinium* cBtQ1 seems to have taken place by the combined effect of the insertion of an IS and a later deletion event, removing the complete *bioB* gene and almost the complete

sequence of the adjacent *bioF* gene (92.5% identical to cEper1 in the remnant segment). Another recent signal of the loss of a nutritional contribution is the pyridoxal-dependent enzyme cystathionine gamma-lyase (involved in the synthesis of cysteine) whose CDS contains an internal stop codon mutation that produces the pseudogene CHV_c0068 in cBtQ1 (94.9% identical to cEper1 gene). A phylogenetic analysis showed that the functional gene, present in this state in cEper1, was acquired by an ancestor through HGT from a eukaryote, perhaps an amoeba (supplementary file S1: supplementary fig. S2, Supplementary Material online). The phylogenetic analysis, including the three in silico identified cystathionine metabolizing enzymes of *L. major* (Williams et al. 2009), showed its closer relation with *L. major* cystathionine gamma-lyase rather than with *L. major* cystathionine beta-lyase, as previously annotated in *Cardinium* cEper1 (Penz et al. 2012).

In *Cardinium* cBtQ1, the inability to synthesize pyridoxal and biotin may be complemented by other facultative endosymbionts of the host. In the case of *B. tabaci* strain QHC-VLC, such enzyme synthesis could be achieved by *Hamiltonella*,

which is located inside the bacteriocytes (supplementary fig. S3, Supplementary Material online). The complete sets of genes required for the synthesis of both cofactors was described for the *Hamiltonella* secondary endosymbiont of the aphid *Acyrtosiphon pisum* (Degnan et al. 2009), and all of them were also found by BLAST similarity in the draft genome of *Hamiltonella* from *B. tabaci* strain QHC-VLC (unpublished data).

Gliding Genes in Cardinium

As stated earlier, the genome of *Cardinium* cBtQ1 harbors four gliding genes organized in an operon (*gldK*, *gldL*, *gldM*, and *gldN*) in the pcBtQ1 plasmid (fig. 3). To state that the differential tissue locations of *Cardinium* endosymbionts of *B. tabaci* (scattered) and *Encarsia* spp. (restricted to ovaries) were correlated with the presence of the four gliding genes, two populations of *E. pergandiella* (USA and Brazil), one of *E. hispida* (Italy) and one from *E. inaron* (USA) were screened for their presence. None of these species gave a positive result for any of the gliding genes, whereas the presence of *Cardinium* was stated in all of them through the amplification of the control gene *gyrB* (supplementary file S1: supplementary materials and methods and tables S4 and S6, Supplementary Material online).

For *B. tabaci*, adult whiteflies were sampled in 12 points from four localities of the province of Valencia (Spain) (supplementary file S1: supplementary materials and methods and tables S5 and S6, Supplementary Material online). Polymerase chain reaction (PCR) amplification revealed the presence of the four gliding genes (as well as of the large plasmid gene *CHV_p021*) in all the samples (except one that failed in all the PCR amplifications). To determine insect biotypes, a mitochondrial cytochrome oxidase I (COI) fragment was sequenced in four females per sampling point. Nine of the samples belonged to the biotype Q (Mediterranean species), whereas two were from the biotype S (Sub-Saharan Africa species), an uncommon biotype in Spain (Moya et al. 2001) (supplementary file S1: supplementary fig. S4, Supplementary Material online). All the samples, including those of biotype S, harbored *Cardinium*, which was determined by PCR amplification of a fragment of the 16S rRNA gene. PCR products from biotype S (sample F) and biotype Q (sample B) (1,100 bp) were sequenced and resulted 100% identical to *Cardinium* cBtQ1.

The presence of additional secondary endosymbionts was also checked finding that biotype Q individuals also contained *Hamiltonella*, whereas biotype S contained *Arsenophonus* and *Wolbachia* (supplementary file S1: supplementary materials and methods and tables S5 and S6, Supplementary Material online).

Discussion

Although the cost of maintaining an obligate mutualistic endosymbiont may be compensated in many cases by its supplementation of the host diet, the maintenance of stable associations with facultative symbionts may require either reproductive manipulation or compensatory benefits from the endosymbiont (Oliver et al. 2010; White et al. 2011; Wernegreen 2012). Both facultative and obligate endosymbionts are transmitted vertically, but facultative symbionts may retain the ability to be horizontally transmitted as revealed by the incongruence of host and symbiont phylogenetic reconstructions (Russell et al. 2003).

Symbionts belonging to the genus *Cardinium* are present in many types of arthropods including arachnids (Nakamura et al. 2009), crustaceans (Edlund et al. 2012), and insects (Zchori-Fein et al. 2004; Zchori-Fein and Perlman 2004; Gruwell et al. 2009; Nakamura et al. 2009), indicating that members of this genus can colonize new niches. Phylogenies and nucleotide conservation of two genes (16S rRNA and *gyrB*) show that *Cardinium* from *B. tabaci* QHC-VLC was closely related to *Cardinium* from armored scale insects and parasitoid wasps, including the species type *C. hertigii* from the wasp *E. hispida* (Zchori-Fein et al. 2004). Researchers have also proposed that *Cardinium* cBtQ1 was a strain of *C. hertigii*, closely related to *Cardinium* cEper1 (symbiont of *E. pergandiella*) whose genome has recently been reported (Penz et al. 2012). In this work, we have sequenced the genome of *Cardinium* cBtQ1, and all the analyses carried out confirm that this strain is closely related to *Cardinium* cEper1, despite being endosymbionts of whiteflies and parasitoids, respectively. There are, however, some differences that could be related to the massive presence of IS in *Cardinium* cBtQ1, as well as to the adaptation to the specific hosts of each strain (see later).

The high number of available Bacteroidetes genomes provided a robust phylogeny (fig. 1), which showed that both *Cardinium* (cEper1 and cBtQ1) and *A. asiaticus* formed a well-defined clade, distant from other family members of Cyclobacteriaceae, Cytophagaceae, and Flammeovirgaceae in the order Cytophagales. Thus, we propose that they form a new family, to be named Amoebophilaceae. This name is proposed because *A. asiaticus* has a larger genome than *Cardinium* spp. and shares more genes with LCA2 than either *Cardinium* strain. Also on the basis of this phylogeny, we were able to infer the coding gene contents (clusters of genes) of the most recent common ancestor in the *Cardinium*/*Amoebophilus* clade (LCA4, LCA2, and LCA1) and to analyze the evolution of the gene repertoires (figs. 6 and 7).

Despite it is not the objective of this work, we can extract general ideas from the cluster analysis (fig. 6). The distribution of genes in COG categories is a well predictor of the way of life of the organisms. Hierarchical clustering indicates that LCA3 to 9 were, similar to free-living Bacteroidetes, able to

occupy different niches (fig. 6). For example, the differences between the abundance of G category in the middle and right clusters could be related to a more restricted source of carbohydrates (niche specialization). It also seems that the increase of the coenzyme metabolism (H) in the middle cluster could be advantageous for the establishment of symbiotic relationships (*Cyclobacterium* was found in the celomic fluid of a sand dollar, [supplementary table S3, Supplementary Material online](#)).

The number of gene clusters estimated in LCA4 was between 1,301 and 1,985, showing a COG profile similar to those of *M. tractuosa* (Flammeovirgaceae) and *Cy. marinum* (Cyclobacteriaceae) (fig. 6), and differing from the COG profiles of LCA2 and LCA1, which clustered with those of the symbiotic bacteria *A. asiaticus* and *Cardinium* (fig. 6). Because species of the family Cyclobacteriaceae are mostly marine free-living bacteria, and associations with animal hosts have been described ([supplementary table S3, Supplementary Material online](#)), we propose that LCA4 was probably a marine free-living bacterium, with a wide range of functional capabilities and the ability to establish symbiotic relationships. Also, it is likely that LCA4 was able to glide because it contained the whole set of gliding genes essential for gliding, including the *sprATE* genes (McBride and Zhu 2013).

The transition from LCA4 to LCA2 was clearly a reductive process that affected almost all COG categories (fig. 7A) producing an ancestral endosymbiont with few biosynthetic capabilities. Considering that the species derived from LCA2 were endosymbionts of amoebas (Horn et al. 2001; Schmitz-Esser et al. 2010) or insects (Zchori-Fein and Perlman 2004; Penz et al. 2012), the most probable reason for this reduction was the transition from a free living to intracellular life style, to start a symbiotic (either mutualistic or parasitic) relationship with a eukaryotic host. During this transition, the number of gene clusters and associated functions was reduced, although LCA2 maintained the ability to acquire new genes by HGT. The higher number of gene clusters in *A. asiaticus* versus LCA2 (298) could be due to this fact, although other reasons, such as the possibility of a biased sample of genomes, or different annotation problems, would also explain its high number of specific gene clusters.

Genome reduction was an ongoing process in the LCA1/*Cardinium* clade, and it was notorious for some categories such as E and G (fig. 7B). Moreover, the comparative analysis of the gene contents of the two *Cardinium* strains revealed that, despite being very similar at nucleotide level (92.9% nucleotide identity), revealing a recent evolutionary divergence, there are some relevant differences between the genomes of both strains, indicating differences in the evolution of endosymbiosis in *Encarsia* spp. and *B. tabaci*.

First, both *Cardinium* contain a plasmid of 50–60 kb with many differences in gene content (fig. 3). However, both plasmids contain a short syntenic block of genes, whose nucleotide content and gene order conservation, as well as the gene

phylogenies carried out, suggests that both derived from an ancestral plasmid present in their LCA (LCA1), and the differences must have been accumulated after the split of both lineages. These differences are due to the insertion of mobile elements, sometimes carrying accessory genes, and to the transfer of genes from the chromosome.

Second, there are also differences in the chromosome of both *Cardinium* strains. The *Cardinium* cBtQ1 chromosome is 126 kb larger (probably a bit larger because, as a draft genome, gaps are not taken into account for this calculation) than that of *Cardinium* cEper1; nevertheless, this difference is not associated with a greater number of genes but to the presence of a large number of pseudogenes, most of them due to defective transposase encoding genes (table 1). The number of repeated sequences in the genome of *Cardinium* cBtQ1 is twice that of *Cardinium* cEper1, and most of these differences are due to a large number of IS in the former genome (fig. 5B). Moreover, some IS types seem to have their origins in Alphaproteobacteria, probably related to the genera *Rickettsia* or *Wolbachia*, supporting the idea of HGT events between secondary endosymbionts harbored by the same host (Toft and Andersson 2010; Schmitz-Esser et al. 2011; Duron 2013). A large number of mobile elements is a typical feature of endosymbionts that have established a recent relationship with their hosts, such as *Candidatus* *Sodalis pierantonius* str. SOPE (Gil et al. 2008, Oakeson et al. 2014) or *Sodalis glossinidius* (Belda et al. 2010). Also, enrichment in mobile elements seems to be linked to genome plasticity in some facultative symbionts (Gillespie et al. 2012). Several lines of evidence indicate that IS elements ([supplementary table S8, Supplementary Material online](#)) are already active in *Cardinium* cBtQ1. This activity combined with both the high number of copies throughout the genome, and a complete replication and repair machinery that can produce recombination, probably underlies the massive number of rearrangements in the genome of *Cardinium* cBtQ1, compared with *Cardinium* cEper1 and *A. asiaticus* (fig. 5C).

Finally, among the genes with an annotated function and differential presence in the two *Cardinium* strains, we consider some may give clues about the relationship between *Cardinium* cBtQ1 and *B. tabaci*. First, the presence of pseudogenes for *pdxT* (pyridoxal biosynthesis) and *cgl* (cystathionine gamma-lyase) and the loss of two genes of the biotin operon indicate that these genes were present in LCA1. This also indicates that the loss of these functions in *Cardinium* cBtQ1 is associated with settlement in a new environment composed by *B. tabaci* and its facultative symbiont *Hamiltonella defensa*. This symbiont seems to have become established in the populations of *B. tabaci* Q1 from Western Mediterranean based on the detection of frequencies of 100% (or almost 100%) in populations from North Africa and south-western Europe (Gueguen et al. 2010). Second, among 71 gene clusters of *Cardinium* cBtQ1, which are absent in *Cardinium* cEper1, there are up to 14 specific genes encoding ankyrin-domain

proteins, which can interact with the host's machinery, but further studies are needed to understand their functions. Finally, the most interesting genes are those whose expression has been amplified by either gene duplication (*mreB*, *lpxH*, *yitW*, and *rtxBDE/toIC*) or by their presence in a multicopy plasmid (e.g., *CHV_p018*, a putative toxin-related gene, *CHV_p021*, a RHS-domain protein, and the gliding genes *gldKLMN*) (supplementary table S2, Supplementary Material online).

On the basis of the ancestor reconstruction analysis, we can predict that the four gliding genes were present in LCA4, LCA2, and LCA1 and were lost in *Cardinium* cEper1. Although LCA4 conserved full gliding machinery, *sprATE* was lost in LCA2 possibly due to its accommodation to an intracellular environment. Because LCA1 conserved the *gldKLMN* operon, this suggests that *gldKLMN* was lost in the *Cardinium* cEper1 lineage. Also, as in the closest Bacteroidetes genomes, such as *A. asiaticus*, *M. tractuosa*, or *Cy. marinum*, these genes are located in the chromosome, and we can postulate that in *Cardinium* cBtQ1, they have been translocated to a multicopy plasmid conserving the operon order. This supports the importance of these genes for *Cardinium* cBtQ1 and suggests that they may explain why the strain is not confined to a single tissue in *B. tabaci* (Gottlieb et al. 2008) (supplementary fig. S3, Supplementary Material online), in opposition to *Cardinium* cEper1 that is restricted to the ovaries of *Encarsia* (Zchori-Fein et al. 2004; Penz et al. 2012). Moreover, the gene amplification in *Cardinium* cBtQ1 not only of the four gliding genes but also of *mreB* and of the cluster *rtxBDE/toIC* suggests that they may play an important role in this organism, as genome reduction is an ongoing process in this strain. There are two possible hypotheses: 1) those genes are involved in gliding as in other genomes; 2) they are involved in the novel type 9 secretion system (PorSS), which is also associated with the secretion of proteins involved in motility and toxins (Sato et al. 2010; McBride and Zhu 2013).

Different Bacteroidetes possess the ability to move by a gliding mechanism, which is related to the ability to degrade some components present in the environment such as chitin and cellulose (Spormann 1999; McBride 2004; Braun et al. 2005). Several examples of gliding have been reported in species of the class Cytophagia where *C. hertigii* was included (Xie et al. 2007; McBride and Zhu 2013), and the scattered pattern detected for the tissue distribution of *Cardinium* cBtQ1 in *B. tabaci* might be caused by a similar mechanism.

The motor model (or focal adhesion) proposed for gliding in myxobacteria (Spormann 1999; Mignot et al. 2007; Jarrell and McBride 2008; Nakane et al. 2013; Nan et al. 2013) could be extended to Bacteroidetes, including *Cardinium* cBtQ1, although in a more simplified manner. This model considers molecular motors that are associated with cytoskeletal filaments and use proton motive force to transmit force through the cell wall to attached dynamic focal adhesion complexes (adhesins) to the substrate, causing the cell to move forward

(Mignot et al. 2007; Sun et al. 2011; Nan et al. 2013). MreB, the homolog of the eukaryotic actin, has been proposed as the cytoskeletal part of the gliding machinery (Kearns 2007; Mauriello et al. 2010). Also, there may be an association with FtsZ, a protein that is part of the bacterium cytoskeleton and can produce force by itself (Erickson et al. 2010). Linkage between the cytoskeleton and gliding is supported by experimental data with the use of compound A22, which is able to affect the MreB structure and inhibits the gliding motility in *Mycobacteria* (Nan et al. 2011). Eleven genes, detected in most Bacteroidetes, have been defined as the core of the gliding machinery. Four of these genes (*gldB*, *gldD*, *gldH*, and *gldJ*) have unknown function, whereas the remaining seven genes (*gldK*, *gldL*, *gldM*, *gldN*, *sprA*, *sprE*, and *sprT*) also encode the PorSS system (McBride and Zhu 2013).

Cardinium cBtQ1 does not show the complete gliding machinery as it only contains four gliding core genes (*gldKLMN*). Neither homologous nor potential analogous genes of *gldBDHJ* have been detected. The *sprAET* genes are also absent, but their function would potentially be substituted by the cluster *rtxBDE/toIC*, which is also duplicated. RTX secretion system belongs to the T1SS and is able to transport proteins from the cytosol to the extracellular space in a SecYEG independent manner. Also, T1SS is able to secrete many different RTX family proteins and proteins without the C-terminal RTX nonapeptide (Linhartová et al. 2010; Kaur et al. 2012). The RTX system would secrete the adhesins (or other proteins that could interact with the host) across the bacterial membrane. However, we did not detect any orthologs to known adhesin proteins, but proteins with eukaryotic domains, such as ankyrins, TPR or WH2 (found in this strain), may function as adhesins in a multicellular eukaryotic organism. Moreover, *Cardinium* may be able to manipulate the host cytoskeleton to form a scaffold, which could be used by the gliding machinery (Haglund et al. 2010). Gliding seems to be a widespread direct invasion mechanism for different kinds of cells (Sibley et al. 1998; Furusawa et al. 2003; Sibley et al. 2004), thus *Cardinium* cBtQ1 and other Bacteroidetes might use this motor model system to invade new hosts or host tissues. In fact, *Cardinium* endosymbiont of *Ixodes scapularis* has been cultivated on insect cell lines and is capable of infecting new cells, even cell lines from different insect species, when they are added to the culture (Morimoto et al. 2006).

The second hypothesis would consider that the *gldKLMN* operon is not involved in gliding, but it is just required for secretion forming the PorSS system, which was initially described for *Porphyromonas gingivalis* as a novel secretion system with eight proteins involved (PorK, PorL, PorM, PorN, PorT, PorW, Sov, and PorP) (Sato et al. 2010). Putatively orthologous genes in the gliding system for the first seven are *gldK*, *gldL*, *gldM*, *gldN*, *sprT*, *sprE*, and *sprA*. The proposed orthologous gene for *porP* in *F. johnsoniae* was *Fjoh_3477*. A similar gene was not detected in either *A. asiaticus* or *Cardinium*. Proteins secreted by the PorSS systems are adhesins, as well as some enzymes

such as chitinases, and gingipains in *F. johnsoniae* and *P. gingivalis*, respectively. Also, proteins secreted by the PorSS secretion system may contain a conserved C-terminal domain (TIGR4131 and 4183) (Sato et al. 2010; McBride and Zhu 2013). However, there were not proteins of *Cardinium* cBtQ1 with this domain. In the PorSS system, the presence of the protein complex GldKLMN is associated with the generation of the energy required for protein secretion by SprTEA. However, these proteins are not encoded in the genome of *Cardinium* cBtQ1 (they are also absent in *A. asiaticus*), and their substitution by the T1SS (RTX system) seems unlikely because T1SS has its own ATP-binding cassette, making the energy production function of the *gldKLMN* unnecessary. This suggests that the PorSS system does not work in *Cardinium* cBtQ1. Thus, we hypothesized that the retention of the necessary genes for the gliding movement (*gldKLMN*) and further amplification by the translocation to a multicopy plasmid, together with the acquisition of the RTX TISS is associated to the scattered phenotype of this strain and the ability to glide.

The frequent presence of *Cardinium* cBtQ1 in *B. tabaci* biotypes Q and S may be due to its ability to benefit its host (Oliver et al. 2008; Feldhaar 2011; Ferrari and Vavre 2011), possibly related to its motility feature, and/or to the presence of some putative toxin-related genes such as *CHV_p018* (low *e*-value BLAST hit against RTX toxins of *H. defensa* from *A. pisum*) and *CHV_p021*, which could play a role in intercellular competition, intercellular signaling, and insecticidal activity based on the presence of the RHS domain (Koskiniemi et al. 2013). As a mobile endosymbiont, *Cardinium* cBtQ1 may contact a parasitoid directly and secrete insecticidal toxins near it, could invade the parasitoid tissue, and kill it by secreting unknown toxins or by the cytotoxic effect of lipid A in a nonacclimated host (Furusawa et al. 2003; Caspi-Fluger et al. 2011; Rader et al. 2012). Furthermore, other effects increasing host fitness cannot be excluded, like heat-stress resistance or maybe some advantages conferred by the lipoate supplementation (Moran et al. 2008; Moya et al. 2008). However, fitness may only be improved in some environments or climate conditions.

In conclusion, we have reported the genome of *Cardinium* cBtQ1 endosymbiont of *B. tabaci*. Comparative genomics and ancestors' reconstruction of gene content have shed light on the drastic reduction in many functional categories that have taken place since its free-living ancestor up to the present. The loss of several cofactors and amino acid biosynthetic capabilities, retained in its close relative *Cardinium* cEper1, rules out an important role in host nutrition and suggests a relationship with its establishment in *B. tabaci* and its endosymbionts. The genome is still very dynamic, with many active transposable elements and rearrangements. On the basis of genomic data, we propose that *Cardinium* cBtQ1 has retained the minimal gliding core machinery present in its ancestors, which in combination with the acquired RTX secretion system might be used to move inside its host or invade new hosts.

Supplementary Material

Supplementary file S1 is available at *Genome Biology and Evolution* online (<http://www.gbe.oxfordjournals.org/>).

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